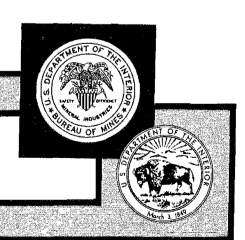
Chromium Recovery From High-Temperature Shift Cr-Fe Catalysts

By A. M. Stubbs and B. W. Jong

BUREAU OF MINES



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UNIT OF MEASURE ABBREVIATIONS USED IN THIS REPORT

degree Celsius lb/d °C pound per day cubic centimeter cm^3 M gal thousand gallon ft foot min minute milliliter mL gram g

gal gallon MMBtu million British thermal units

pct

percent

g/L gram per liter mt metric ton

kW kilowatt ppm part per million

L liter st short ton

1b pound yr year

h

hour

CHROMIUM RECOVERY FROM HIGH-TEMPERATURE SHIFT Cr-Fe WASTE CATALYST

By A. M. Stubbs¹ and B. W. Jong¹

With an Appendix on Process Economics by M. J. Magyar

ABSTRACT

The Bureau of Mines has devised a procedure to selectively extract and recover chromic oxide (Cr₂O₃) from waste high-temperature shift catalysts that contain approximately 6 pct Cr and 62 pct Fe, both in oxide form. The Bureau's procedure consists of roasting the waste catalyst with sodium hydroxide (NaOH) or sodium carbonate (Na₂CO₃) followed by water leaching. The chromium from the leach liquor was recovered utilizing an oxidation-reduction reaction with pH adjustment to form a hydrous chromic oxide (Cr₂O₃•XH₂O) precipitate. The roast-leach procedure produced chromium extractions greater than 89.7 pct, with product recovery in excess of 99 pct from the leach liquor.

The costs of the NaOH and Na₂CO₃ roast processes, including sizing and equipment-related recommendations, were determined by the Bureau's Office of Process Evaluation. The economic evaluation indicates that producing a Cr₂O₃ product, at the present time, is not competitive with the present-day cost of imported chromites. The estimated average operating cost for both processes was \$5.44/lb of recovered Cr₂O₃ concentrate. However, this procedure offers a potential alternative to the disposal of this waste through landfill.

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INTRODUCTION

The United States is heavily dependent upon foreign sources for chromium to support its metallurgical, chemical, and refractory industries (1).² Therefore the Bureau of Mines has investigated several approaches for the recovery of chromium from waste catalysts.

In the production of ammonia, a Cr-Fe containing catalyst is used at high temperatures to increase the hydrogen content of synthesis gas. After a period of time in use, this catalyst becomes contaminated with sulfur. At a predetermined point, the catalyst is discharged from the equipment and is considered a waste product. The waste product is generally disposed through landfill. To date, commercial recycling has not been justified because of transportation and refining costs, but most manufacturers express interest in obtaining an outlet for utilization of the waste catalyst rather than disposal in landfills (1).

There is 24.5 million lb of this type of catalyst in place (1). Annual catalyst discharge is 7 million lb, which amounts to an annual chromium loss of 420,000 lb. As of December 1985, the annual U.S. consumption of chromium was 585 million lb.

Previous Bureau work by Hundley (3), based partially on earlier work by Chandra (4), had demonstrated Cr could be recovered from various types of low-grade chromite ores using molten NaOH under oxidizing conditions. Current industry practices use Na₂CO₃ and a lime (CaO) mixture to recover Cr from high-grade ores (4). At this time, few if any studies have been conducted aimed at recycling the metal content of high-temperature shift catalysts because of their low-chromium content and location (1).

The Bureau's approach to extract and recover Cr from spent high-temperature shift catalyst is shown in figure 1. Briefly, the procedure consisted of mixing the waste catalyst with a NaOH or Na₂CO₃ reagent followed by addition of water. This mixture was then roasted at 600°

to 700° C to form sodium chromate (Na₂CrO₄). The catalyst was also mixed with hydrogen peroxide (H₂O₂), reacted, and then mixed with a NaOH or Na₂CO₃ solution and roasted at 600° to 700° C to form Na₂CrO₄. The basic reactions occurring during roasting of the waste catalyst using NaOH and Na₂CO₃ with or without H₂O₂ follow:

$$Cr_2O_3 + 4NaOH + \frac{3}{2}O_2 \rightarrow 2Na_2CrO_4 + 2H_2O \uparrow$$
 $Cr_2O_3 + 4NaOH + H_2O_2 + O_2 \rightarrow 2Na_2CrO_4 + 3H_2O \uparrow$
 $Cr_2O_3 + 2Na_2CO_3 + \frac{3}{2}O_2 \rightarrow 2Na_2CrO_4 + 2CO_2 \uparrow$
 $Cr_2O_3 + 2Na_2CO_3 + H_2O_2 + O_2 \rightarrow 2Na_2CrO_4$
 $+ H_2O \uparrow + 2CO_2 \uparrow$

The roasted product was then ground and leached with hot water to extract the Cr. The leach liquor was treated by two different methods for Cr recovery. The first method used sodium sulfite (Na₂SO₃) or sodium bisulfite (NaHSO₃) for reduction of Cr to the trivalent form, followed by pH adjustment with sulfuric acid (H₂SO₄) to around 5.0 and NaOH pH adjustment to 7.5 for precipitation. The second method required that the pH be lowered with H₂SO₄, and then hydrazine (N₂H₄) was used as the reductant to adjust the pH. The precipitate produced by both methods, identified by X-ray diffraction, was of the form Cr₂O₃•XH₂O, which agreed with the findings by Udy (5). The product was calcined at 350° C to produce Cr₂O₃. The waste solution produced from the precipitation was purified by stripping the Na⁺ and SO₄² ions with an electrodialysis ion-selective membrane.

This report discusses the experimental results from laboratory-scale extraction and recovery studies of Cr from a Cr-Fe high-temperature shift catalyst. Reported are the optimized parameters for the unit operations of roasting, leaching, and product recovery.

²Italic numbers in parentheses refer to items in the list of references preceding the appendix at the end of this report.

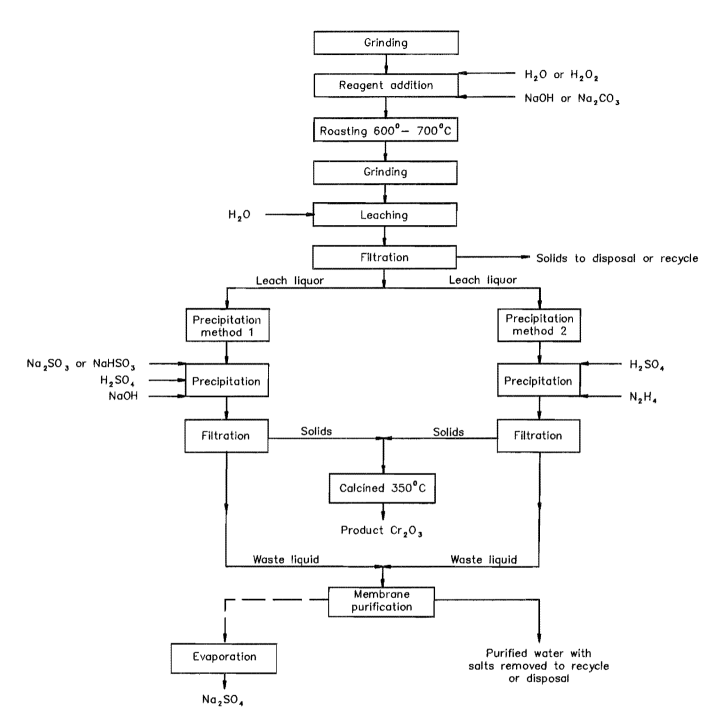


Figure 1.-Process flowsheet for Cr recovery from Cr-Fe waste high-temperature shift catalyst.

RAW MATERIAL

The catalyst studied in this report was obtained from Inco Research & Development Center, Inc., Sterling Forest, Suffen NY through Bureau contract J0215042 (1). No information was obtained or given by Inco disclosing catalyst origin or if the sample was typical.

The catalyst was received in pellet form. The major phases in the pellets contained both Cr₂O₃ and ferric oxide (Fe₂O₃). Chemical analysis of the material showed that the catalyst contained 5.38 pct Cr, 61.3 pct Fe, 0.71 pct Ti, 0.61 pct Mg, 0.30 pct Mn, and 0.23 pct Si.

EQUIPMENT AND EXPERIMENTAL PROCEDURES

The roasting of the waste catalyst was carried out in a small, 2,500-cm³ muffle furnace with a 94-cm³ high-temperature aluminum oxide (Al₂O₃), ceramic crucible. No special mixing or gas injection system was used to enhance the roast or the subsequent water leach. Water leaching of the roasted product was done in a 250-mL, closed, three neck flask with agitation, followed by gravity filtration. Product precipitation was done in an open beaker with heating. Following the precipitation of the Cr₂O₃•xH₂O product, salts in the residual solution were concentrated with an IONICS³ MEDIMAT5 electrodialysis system with a Cr51CZ1386 cation, AR103QZL386 anion membrane to produce water for reuse or discharge.

Chromium extraction tests consisted of placing 6.0 g NaOH or 6.3 g Na₂CO₃ in a high-temperature ceramic crucible with 25.0 g of catalyst, ground to minus 60-mesh. Twenty-three milliliters of water was added and the combination was mixed until the chemical reagent was dissolved. The reaction mixture was then roasted at 600° to 700° C for 2 h.

An alternate experimental method used 23 mL of 8.0 pct H_2O_2 added to 25.0 g of minus 60-mesh catalyst. The mixture was allowed to react 8 to 10 min and NaOH or Na_2CO_3 was then added, 6.0 and 6.3 g respectively, and allowed to dissolve. The mixture was roasted for 2 h at 600° to 700° C.

To determine the amount of Cr extracted from the waste catalyst, the roasted material, a lightly sintered, easily crushed mass, was ground to minus 60 mesh with mortar and pestle. The minus 60-mesh powder was water

leached at 5.0 pct solids for 2 h at 100° C and then gravity filtered. The leach liquor and solids were analyzed for Cr. The extraction percentage represented the amount of Cr in solution, based upon total Cr in the leach solution and remaining residue.

The method for recovering Cr from the leach liquor used either Na₂SO₃ or NaHSO₃ as reducing agents. The reducing agent was added as a powder in appropriate amounts based on 25 g/L of salt. H₂SO₄ (12<u>M</u>) was added to reduce the 12.0 to 12.5 pH level to 3.1 to 5.0 with a pH of 5.0 being preferred to initiate the oxidation-reduction reaction between the Na₂CrO₄, NaHSO₃, or Na₂SO₃, and the H₂SO₄ in solution. Final pH adjustment to 6.8-7.5 was made by adding a 100-g/L-NaOH solution and heating to 60° to 65° C, which yielded the precipitated product.

A second Cr recovery method used the same pH adjustment. The pH level was then increased to 6.8-7.5 with a 35-pct solution of N₂H₄, which also served as the reductant, to form the precipitate. No external heat was necessary with this method.

The Cr recovery by both precipitation techniques was calculated as the amount precipitated from the leach liquor. The Cr precipitate produced by both recovery methods was calcined at 350° C for approximately 10 min to produce a dried Cr₂O₃ product.

The variables studied in this investigation were the reagent-to-catalyst ratio, roast time, roast temperature, leach temperature, leach water volume, and the amounts of reductants used for chromium precipitation.

RESULTS AND DISCUSSION

CATALYST ROAST

The Cr-Fe waste catalyst (25.0 g) was initially roasted for 2 h using a 4:1 catalyst-to-Na₂CO₃ ratio. The roast was conducted using temperatures ranging from 500° to 900° C with the initial roast material in dry, paste, or wet form depending on test conditions. Results are shown in figure 2. Paste roasting with 15.0 mL of water produced the

³References to specific products does not imply endorsement by the Bureau of Mines.

highest extraction values at temperatures between 700° and 900° C. Extraction values of paste and dry roasting were similar at 600° C. Wet roasting with 30.0 mL of water showed poor extraction values at all temperatures, owing primarily to excessive spattering, and was not considered further. Dry roasting with no addition of water resulted in 5.0 to 6.0 pct lower extraction values than paste roasting at 700° C. Higher dry roasting temperatures to increase extractions were not attempted, since extrapolation of the dry roast data (fig. 2) indicated no significant advantage would likely result.

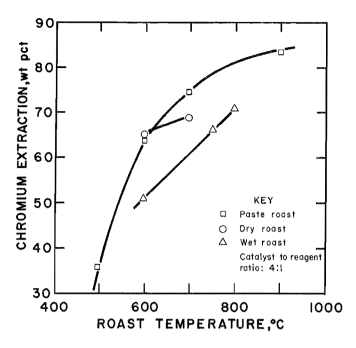


Figure 2.—Effect of roast temperature and material condition of Cr extraction.

An attempt was made at increasing the extraction efficiency by increasing the roast time of the dry roast from 2 h to 3 and 4 h at 600° and 700° C. The test results indicated (table 1) that extractions increased at 600° C giving comparable results to paste mixing at 700° C for 2 h (fig. 2), with an average extraction of 75.0 pct. The 700° C temperature produced extractions 3 to 4 pct lower than 600° C. This result is possibly due to carbonate loss at 700° C, which is 23.0 pct greater than at 600° C as shown in table 1. Dry roasting was discontinued, since paste roasting gave relatively the same extraction values with shorter roast times. Paste roasting at 700° C for 2 h was used as the standard roasting technique for further testing.

NaOH was substituted for Na₂CO₃ at a weight ratio of 4.2:1 and paste roasted for 2 h at 700° C. The Cr extraction was 76.9 pct, which was comparable to the Na₂CO₃ approach at 700° C as shown in figure 2.

The amount of water used to mix the catalyst and NaOH was increased from 15.0 to 23 mL, to increase reagent solubility; roast time and temperature remained constant. The metal extraction increased only from 76.9 to 78.9 pct.

Tests were run using catalyst-to-reagent weight ratios of 8.3:1, 4.2:1, 2.8:1, and 2.1:1 at 700° C for 2 h and using

TABLE 1. - Chromium extraction by dry roasting

Roast time, h	Roast temperature, °C	Extraction pct	Carbonate loss, pct
3	500	76.3	58.3
4	600	74.6	54.7
3	700	72.0	81.7
4	700	71 <u>.5</u>	76.9

23 mL water with both NaOH and Na₂CO₃. The results are listed in table 2. Chromium extraction increased with decreasing ratio to 2.8:1 for NaOH (89.7 pct) and 2.1:1 for Na₂CO₃ (90.9 pct). Results from NaOH tests showed an extraction decrease at a ratio of 2.1:1. Lowering the catalyst-to-NaOH ratio to 1.3:1 resulted in an extraction decrease to 47.0 pct.

H₂O₂ was substituted for water in various concentrations ranging from 2.0 to 30.0 pct as an oxidizing agent, while maintaining previous experimental conditions at 700° C for 2 h and 23 mL of H₂O₂ solution. Since an oxidizer was used, it was decided to maintain the amount of NaOH and Na₂CO₃ used initially; i.e., 6.0 g NaOH and 6.3 g Na₂CO₃. The resulting Cr extraction utilizing NaOH and Na₂CO₃ reagents ranged from 84.0 to 93.0 pct. NaOH produced 1.0 to 6.0 pct greater extractions than Na₂CO₃ with the same H₂O₂ addition (table 3).

Tests were run for both NaOH and Na₂CO₃ at various catalyst-to-reagent ratios with 8.0 pct H₂O₂ solution. The Cr extractions were higher than those obtained when water was used in the paste. The use of H₂O₂ during the roast preparation increases extractions around 5.9 pct for NaOH solutions and 5.6 pct for Na₂CO₃ solutions as shown in tables 2 and 4 and figure 3.

LEACHING

Twenty-five grams of roasted product was initially leached in water at 5.0 pct solids, ambient pressure, and 100° C for 2 h to extract Cr as Na₂CrO₄. The Cr⁶⁺ concentration produced under these conditions was 6.0 g/L with impurity levels of <0.1 ppm Mg and Mn, 7.5 ppm Ti, and 25 ppm Si. The impurity concentrations remained constant under extraction studies carried on

TABLE 2. - Chromium extraction for various water-mixed catalyst-to-roast reagent ratios, percent

	Catalyst-to-reagent ratio	NaOH	Na ₂ CO ₃
8.3:1	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	65.7	48.1
4.2:1		83.7	73.0
		89.7	80.6
		87.9	90.9

TABLE 3. - Chromium extraction using various amounts of H_2O_{21} percent

(700° C for 2 with 4.2:1 catalyst-to-reagent ratio)

H_2O_2 , pct	NaOH	Na ₂ CO ₃
2.0	91.4	86.1
4.0	91.0	84.8
8.0	91.0	87.6
15.0	89.8	88.5
30.0	92.9	91.3

TABLE 4. - Chromium extraction for various H₂O₂-mixed catalyst-to-roast reagent ratios, percent

-	Catalyst-to-reagent ratio	NaOH	Na ₂ CO ₃	
8.3:1		72.2	49.7	
		91.0	87.6	
		96.2	84.0	
		85.9	93.7	

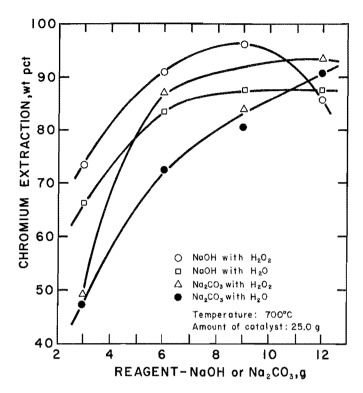


Figure 3.—Study of Cr extraction with varying reagent amount.

later. Extraction studies were made by varying the amount of water used, leach temperature, and leach time. Results are shown in table 5. Based on the extraction studies, the leaching conditions selected were 15.0 min of leach time, a leach slurry of 16 pct solids, and a temperature of 80° C to minimize evaporation from the solution while maintaining a high extraction. All tests were done with NaOH roasted product. These same preferred leaching conditions were applied to Na₂CO₃ roast product (fig. 3).

CHROMIUM PRECIPITATION

Cr₂O₃ was chosen as the recovery product from the leach liquor. This decision was based partly on private communications with the officials in the waste catalyst regeneration industry. It was implied that the catalyst

TABLE 5. - Studies of atmospheric leach testing using NaOH solution

Varied leach slurry 100° C and 2 h		Varied leach temp at 16 pct solids and 2 h		Varied leach time at 16 pct solids and 100° C	
Solid,	Extraction,1	Temp, °C	Extraction,1	Time, h	Extraction,1
5.0	91.3	21.5	71.3	0.25	87.5
10.0	86.4	40.0	72.6	1.0	87.5
16.0	91.3	68.0	74.7	1.5	87.5
25.0	72.8	80.0	85.0	2.5	87.5
		100.0	91.3	I	

¹Values were from stage leach testing and represent amount of Cr⁶⁺ in solution.

manufacturers, which often retain rights over the waste catalysts, would likely require the metal recovered from catalysts to be returned. The decision to recover a Cr₂O₃ product was also based on the relative simplicity of this approach.

The chromate ion was precipitated from the leach liquor through reduction to a chromic ion and then pH adjustment. Na₂SO₃ or NaHSO₃ was added to the leach liquors in concentrations ranging from 10.0 to 60.0 g/L. The pH was lowered to 5.0 by addition of a concentrated H₂SO₄ to allow reduction of Cr⁶⁺ to Cr³⁺. The solution was then heated to 63° or 68° C. The pH was then readjusted with a NaOH solution to a level between 6.8 and 8.1. The Cr recovery from the leach liquors treated with Na₂SO₃ was greater than 99.0 pct (table 6). Optimization of Cr recovery with NaHSO3 was not done since the SO₃² was oxidized in both Na₂SO₃ and NaHSO₃ to SO₄². Product purity results indicate comparable products when either reducing agent was used (table 7). A pH adjustment to lower than 4.5 will produce unwanted hydroxides and/or sulfates.

Use of N₂H₄ as a reducing agent for the chromate ion in leach liquors also was evaluated. A leach liquor volume of 0.15 L was mixed with various volumes of N₂H₄. Addition of N₂H₄ to leach liquors increased the temperature from ambient to 35° to 40° C. N₂H₄ was added at pH 5.0, which reduced the chromate ion from Cr⁶⁺ to Cr³⁺, and resulted in a pH adjustment to a level between 6.8 and 7.4. The solutions were not heated as was the case with either Na₂SO₃ or NaHSO₃. Use of N₂H₄ produces a comparable Cr recovery to the sulfites (99.0 pct or greater) as shown in table 6.

TABLE 6. - Studies of Cr recovery using Na₂SO₃ and N₂H₄ (0.15 L solution)

	NaOH		Na ₂	CO ₃
	Temp,	Recovery, pct	Temp, °C	Recovery,
Na ₂ SO ₃ , g/L: 60				
60	63	99.4	68	99.9
25	63	100	68	100
20	63	82.9	68	99.4
10	63	51.0	68	55.1
N ₂ H ₄ , mL:				
N ₂ H ₄ , mL: 1.3	NAp	NAp	35	53.7
3.3	35	99.0	35	100
4.6	35	100	NAp	NAp
6.6	40	100	35	100
10	35	99.8	NAp	NAp

TABLE 7. - Purity of final Cr₂O₃ products, percent

From NaOH solution using N ₂ H ₄		From NaOH solution using NaHSO ₃		From Na ₂ CO ₃ solution using Na ₂ SO ₃	
Compo- sition	Product	Compo- sition	Product	Compo- sition	Product
Cr ₂ O ₃	94.66	Cr ₂ O ₃	94.58	Cr ₂ O ₃	92.45
Na ⁺ SO ₄ ²⁺	3.09 2.25	Na ⁺ SO ₄ ²⁺	1,39 4.03	Na ⁺ SO ₄ ²⁺	5.45 1.94

The final product, identified as $Cr_2O_3 \cdot XH_2O$, was obtained by calcining the precipitated $Cr_2O_3 \cdot XH_2O$ at 350° C for 5 to 10 min. The purity of the calcined Cr_2O_3 product from NaOH leach solution using N_2H_4 or NaHSO₃ and the Cr_2O_3 product from Na_2CO_3 leach solution using Na_2SO_3 is shown in table 7. Each method produced a product of 92 pct purity or greater.

WASTE WATER TREATMENT

The waste water from the precipitation contained large quantities of Na⁺ and SO₄²⁻ ions with less than 0.1 ppm Cr⁶⁺. A membrane purification method, electrodialysis, stripped the ions and effectively purified the waste water. On the average, the procedure showed that about 70 pct of the waste water was suitable for recycling, the remaining 30 pct required further processing.

Ninety-eight pct of the Na⁺ and SO₄² ions contains in the waste solution from the Na₂CO₃ process were removed. Solutions from the NaOH process produced similar results of 99.2 pct removal. Waste treatment data are listed in table 8. Results show that the purified water

TABLE 8. - Results of waste water treatment, grams per liter

Salt	Feed	W	ater
		Purified	Waste
	NaOH WAST	E SOLUTION	
Na ⁺	4.80	0.02	16.60
Na [†] SO ₄ ²	11.7	.05	8.38
Cr ⁶⁺	≤.0001	≤.0000312	≤.0000688
	Va₂CO₃ WAS1	E SOLUTION	
Na ⁺	3.80	0.07	13.30
Na ⁺	8.44	.13	33.80
CO ₃ ²	.10	.06	.14
<u>Cr⁶⁺</u>	.0002	.0000591	.000141

containing less than 0.07 g/L Na⁺, 0.13 g/L SO₄², 0.06 g/L CO₃², and 0.03 ppm Cr⁶⁺ can be recycled and used in the leaching circuit.

The waste water from the membrane system, which averaged 30 pct of the total sent to the electrodialysis unit, contained the concentrated Na⁺ and SO₄² salts in solution. The solution at this point could possibly be evaporated to recover sodium sulfate (Na₂SO₄), but this was never actually attempted.

PROCESS ECONOMICS

A preliminary economic evaluation on the proposed processes to extract and recover Cr from spent Cr-Fe waste high-temperature shift catalyst was prepared by the Bureau's Office of Process Evaluation.

The estimated capital costs of the plants, each designed to process 9,600 lb of the spent Cr-Fe catalysts, are \$3.9 million and \$4.2 million for the NaOH and the Na₂CO₃ processes. Total operating costs were estimated to be \$5.41/lb and \$5.48/lb of recovered Cr₂O₃ concentrate, for each process, respectively. The costs of the two processes are very close and can be considered to be same because of the estimating technique used for some of the equipment sizes and costs. High-cost items include raw materials at \$0.65/lb of product produced for the NaOH process and \$0.44/lb for the Na₂CO₃ process. Both processes have high labor related charges of approximately \$2.00/lb of product, which include direct labor, plant maintenance, payroll overhead, and operating supplies.

Imported Turkish Cr₂O₃ ore (48 pct Cr₂O₃) costs \$125/mt f.o.b. and adjusting to 99 pct Cr₂O₃, yields a value

of \$258/mt or \$0.13/lb of Cr₂O₃. The currently designed catalyst plants, which are not optimum plant designs because of the limited amount of material, are uneconomical with operating costs of \$5.41/lb and \$5.48/lb of Cr₂O₃ versus a Cr₂O₃ value of \$0.13/lb.

However, the economic situation could improve if alternative Cr_2O_3 and Fe_2O_3 products are produced. An example of this would be to produce high-purity, pigment-grade Cr_2O_3 and Fe_2O_3 . Pure Cr_2O_3 and pure Fe_2O_3 sell at \$1.90/lb and \$0.30/lb, respectively, as pigments. Because of its larger quantity of iron (60 pct Fe versus 6 pct Cr in the waste Cr-Fe catalyst), the Fe_2O_3 filter cake would have a value of \$2.93/lb of Cr_2O_3 produced. Total value of the product would be \$4.83/lb of Cr_2O_3 produced. The original object of this project was to recover Cr and not Fe. There is no information at this time on whether the necessary purity levels are attainable or what additional processing steps would be needed. Any further processing would increase both the capital and operating costs proportionately.

CONCLUSIONS

Laboratory-scale testing was conducted by the Bureau to extract and recover Cr from waste Cr-Fe catalysts. The extraction procedure required that the catalyst be roasted with NaOH or Na₂CO₃ followed by water leaching. The Cr extraction as Na₂CrO₄ was greater than 89.7 pct. Recovery of as Cr₂O₃ was through a reduction, a precipitation, and a calcination step that produced

recoveries of greater than 99 pct, generally leaving <0.02 ppm Cr in solution.

Cost studies were made on both NaOH and Na₂CO₃ roast processes by the Bureau's Office of Process Evaluation. Based on the cost evaluation data, the plant would operate at 9,600 lb/d of waste catalyst processed for both processes. The total operating costs of the two

processes were found to be \$5.41/lb and \$5.48/lb of recovered Cr₂O₃ product for each process respectively. The operating costs of the two processes are very close

with neither process offering a significant cost advantage. However, this procedure offers a potential alternative to the disposal of this waste through landfills.

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APPENDIX.-PROCESS ECONOMICS

By M. J. Magyar¹

Capital and operating costs for two plants are based on the process flowhseet presented in the main text. Each plant is designed to process 9,600 lb/d of spent Cr-Fe high-temperature shift catalysts to recover a Cr₂O₃ product. Major items of equipment for both processes used for cost evaluation are shown in table A-1. The major difference between the two processes is material handling equipment necessary for the NaOH reagent (e.g., unloading, storage). Byproduct recovery was not included.

CAPITAL COSTS

The capital costs estimate is of the general type called a study estimate by Weaver (6).² This type of cost estimate is usually expected to be within 30 pct of the cost for the plant described. However, recent studies on firstof-kind plants show that this kind of accuracy prediction is Because of the limited engineering data deceptive. available and the lack of impurity information, no accuracy limits for this evaluation are predicted.

The estimated fixed capital costs are \$3.6 million and \$3.9 million based on a first quarter 1987 basis (Marshall and Swift (M and S) index of 803.7) for plants processing 9,600 lb/d of spent Cr-Fe high-temperature shift catalysts. Capital costs are listed in Table A-2 for the two plants based on either a NaOH or a Na₂CO₃ roast process. The plants operate 1 shift per day, 5 days per week, 330 days per year except for the roasting and product recovery sections, which operate 3 shifts per day 7 days per week. The remaining 35 days are for scheduled and unscheduled downtime.

Equipment costs used in this estimate are based on informal cost quotations from equipment manufacturers and on capacity cost data. In developing the plant capital costs, corrosion-resistant materials of construction are used where appropriate,

TABLE A-1. - Major items of equipment

Section and item	Quantity	Unit size
Reagent-feed preparation: Roll		
crusher	1	1 st/h
Roasting:		,
Rotary kiln	1	2-ft diam by 21 ft
Rotary cooler	1	1.5-ft diam by 11 ft
Roll crusher	1	1 st/h
Leaching: Leach tank	1	2,900 gal
Precipitation: Precipitation tank .	1	2,000 gal
Product recovery:		,
Rotary dryer	1	2.7-ft diam by 44 ft
Membrane cell	1	400 cell pairs

Working capital is defined as the funds in addition to fixed capital, land investment, and startup costs that must be provided to operate the plant. Working capital, also shown in table A-2, is estimated from the following items: (1) raw materials and supplies inventory (costs of raw material and operating supplies for 30 days); (2) product and in-process inventory (total operating cost for 30 days); (3) accounts receivable (total operating cost for 30 days); (4) available cash (direct expenses for 30 days).

Startup costs are estimated as 10 pct of the fixed capital costs of which 1 pct is shown in table A-2. The remaining 9 pct is assumed to be first year operating costs; however, they are not shown in the operating cost table, but rather are used in calculating the required selling price. Land investment is not included in this estimate.

OPERATING COSTS

The estimated annual operating costs for NaOH and Na₂CO₂ processes are approximately \$1.3 million each as shown in Table A-3. An average of 330 days of operation per year is assumed over the life of the plants. Based on processing 9,600 lb/d of spent catalysts, this corresponds to an annual operating cost of \$0.41/lb of catalyst with an average cost of \$5.44/lb for each process. Operating costs are divided into direct, indirect, and fixed costs.

TABLE A-2.- Estimated capital costs: NaOH roast process versus Na₂CO₃ roast process

	NaOH	Na ₂ CO ₃
	roast	roast 🏻
Fixed capital:		
Reagent-feed preparation section	\$526,900	\$771,000
Roasting section	729,200	722,000
Leaching section	500,100	500,100
Precipitation section	558,100	564,300
Product recovery section	462,100	455,000
Steamplant	39,100	39,100
Subtotal	2,815,500	3,051,500
Plant facilities, 10 pct of above		
subtotal	281,600	305,200
Plant utilities, 12 pct of above		
subtotal	337,900	366,200
Total plant cost	3,435,000	3,722,900
Land cost	0	0
Subtotal	3,435,000	3,722,900
Interest during construction period	143,100	155,200
Fixed capital cost	3,578,100	3,878,100
Working capital:		
Raw material and supplies	15,200	11,300
Product and in-process inventory	105,600	106,900
Accounts receivable	105,600	106,900
Available cash	59,500	57,500
Working capital cost	285,900	282,600
Capitalized startup costs	35,800	38,800
Total capital cost	3,899,800	4,199,500
1		

¹Basis: M and S equipment cost index of 803.7.

¹Chemical engineer, Office of Process Evaluation, Bureau of Mines,

Washington, DC.

Italic numbers in parentheses refer to items in the list of references preceding the appendix.

Direct costs include raw materials, utilities, direct labor, plant maintenance, payroll overhead, and operating supplies. Raw materials and utility requirements per pound of Cr₂O₃ are shown in table A-3. The direct labor cost is estimated on the basis of assigning 4.2 employees for each position that operates 24 h/d, 7 days per week and 1.0 employee for each position that operates 8 h/d, 5 days per week. Payroll overhead includes vacation, sick leave, social security, and fringe benefits.

Indirect costs include the expenses of control laboratories, accounting, plant protection and safety, plant administration, marketing, and company overhead. Research and overall company administrative costs outside the plant are not included.

Fixed costs include the cost of taxes (excluding income taxes), insurance, and depreciation. Depreciation is based on a straight-line, 10-yr period.

TABLE A-3. - Estimated annual operating costs: NaOH roast process versus Na₂CO₃

	NaOl	ł roast	Na ₂ C	CO ₃ roast
	Annual cost	Cost/ib Cr ₂ O ₃	Annual cost	Cost/lb Cr ₂ O ₃
Direct cost:				
Raw materials:				
Na ₂ CO ₃ at \$125.00 per ton	NAρ	NAp	\$78,700	\$0.331
NaOH at \$175.00 per ton	\$144,400	\$0,608	14,400	.061
H₂SO₄ at \$48.00 per ton	8,700	.037	10,100	.043
$N\tilde{a}_{2}S\tilde{O}_{3}$ at \$0.24	100	.001	100	.001
Chemicals for steamplant water treatment	200	.001	200	.001
Total	153,400	.647	103,500	.437
Utilities:	-			
Electric power at \$0.05/kW•h	53,600	.226	60,100	.253
Process water at \$0.20/Mgal	400	,002	400	.002
Natural gas at \$3.00/MMBtu	<u>58,</u> 100	,245	58,100	.245
Total	112,100	.473	118,600	.500
Direct labor:				
Labor at \$10.50/h	152,900	.644	152,900	.644
Supervision, 15 pct of labor	22,900	.096	22,900	.096
Total	175,800	.740	175,800	.740
Plant maintenance:				
Labor	72,300	.304	78,400	.330
Supervision, 20 pct of maintenance labor	14,500	.061	15,700	.066
Materials	72,400	.305	78,300	.330
Total	159,200	.670	172,400	.726
Payroll overhead, 35 pct of above payroll	91,900	.387	94,500	.398
Operating supplies, 20 pct of plant maintenance .	31,800	.134	34,500	.145
Total direct cost	724,200	3.051	699,300	2.946
Indirect cost, 40 pct of direct labor and maintenance	134,000	.564	139,300	.586
Fixed cost:				
Taxes, 1.0 pct of total plant cost	34,400	.145	37,200	.157
Insurance, 1.0 pct of total plant cost	34,400	.145	37,200	.157
Depreciation, 10-yr life	_357,800	1.506	387,800	1.632
Total operating cost	1,284,800	5.411	1,300,800	5.478